## **Attachment 2**

## Clean version of the specification with changes incorporated

1. On page 23, the 1<sup>st</sup> full paragraph starting on line 13 should read as follows:

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The homozygous ixr1-1 (SEQ ID NO. 1) and ixr1-2 (SEQ ID NO. 2) mutant plants are 300 and 90 times respectively, more resistant to isoxaben than wild type plants (Heim et al. 1989). ixr1-1 (ixr1-2 was not tested) is also more resistant to a new thiazolidinone herbicide (TZ, compound 1). At 12µM the thiazolidinone herbicide kills wild type Arabidopsis, but reduces growth of the ixr1-1 mutant by only 50% (Sharples et al., 1998)

- 2. On page 17, the 1st paragraph starting on line 1 should read as follows:

  oligonucleotide primers (F: 5' CGAACTTGAGACCTCTTGA 3 ') (SEQ ID NO. 4); ( R: 5'
  GCTTACCTGGAGACAGTCA 3') (SEQ ID NO. 5) were designed with the Oligo Primer Analysis
  Software version 5.0 (National Biosciences, Plymouth, MN) to PCR-amplify a 124 bp fragment
  from genomic wild-type Columbia DNA, containing the (TG)<sub>13</sub> dinucleotide repeat. Using the
  same primers a shorter PCR-product (<120bp) is obtained from wild-type Landsberg genomic
  DNA, whereas a Col x Ler heterozygous plant give both products, as can be easily seen on 4%
  agarose-gels. PCR-conditions for SSLP-marker med24.2: 50 mM KCL, 10mM Tris-HCL (pH
  9.0 @ 25°C), 0.1% Triton X-100, 200μM dATP, dGTP, dTTP, dCTP (each), 5 pmoles primer F,
  5pmoles primer R, 2.0mM MgCl<sub>2</sub>, 1.0 Units *Taq*. Polymerase (Promega, Madison, WI), 10-50 ng
  genomic DNA, final volume 20μl. PCR-program: 1 min 94°C; 40 cycles (20 sec 94°Cm 20 sec
  55°C, 40 sec 72°C), 3 min 72°C.
- 3. On page 17, the 1<sup>st</sup> full paragraph starting on line12 should read as follow:

  <u>SSLP-marker moj9.2</u> (approx. 100kb proximal of m217 on the physical map of chromosome V; <a href="http://www.kazusa.or.jp/arabi/chr5/pmap/P1">http://www.kazusa.or.jp/arabi/chr5/pmap/P1</a> map 1.html)

molecular target:

(TA)<sub>19</sub> dinucleotide repeat (bases 53618-53655) on chromosome V BAC-

clone MOJ9

PCR primers:

F: 5' CATGATCCATCGTCTTAGT 3' (SEQ ID NO. 6)

R: 5' AATATCGCTTGTTTTTGC 3' (SEQ ID NO. 7)

PCR-product size:

179 bp in Col, ca. 160b bp in Ler

PCR-conditions:

as for med24.2 with 2.2mM MgCl<sub>2</sub>

4. On page 18, the 1<sup>st</sup> full paragraph starting on line 5 should read as follows:

SSLP-marker muk11.1 (approx. 500 kb proximal of med24.2 on the physical map of chromosome V; <a href="http://www.kazusa.or.jp/arabi/chr5/pmap/P1">http://www.kazusa.or.jp/arabi/chr5/pmap/P1</a> map 1.html)

molecular target:

(GA)<sub>38</sub> dinucleotide repeat (bases 57187-57252) on chromosome V BAC-

clone MUK11

PCR primers:

F: 5' TCCAAAGCTAAATCGCTAT 3' (SEQ ID NO. 8)

R: 5' CTCCGTCTATTCAAGATGC 3' (SEQ ID NO. 9)

PCR-product size:

177 bp in Col, ca. 120b bp in Ler

PCR-conditions:

as for med24.2.

5. On page 18, the 2<sup>nd</sup> full paragraph starting on line 12 should read as follows:

SSLP-marker nga158 (approx. 500kb distal of moj9.2 on the physical map of chromosome V; http://www.kazusa.or.jp/arabi/chr5/pmap/P1\_map\_1.html)

molecular target:

(CT)<sub>14</sub> dinucleotide repeat (bases 19384-19411) on BAC-clone MJJ3

PCR primers:

(http://www.kazusa.or.jp/arabi/chr5/map/0-2Mb.html)

F: 5' ACCTGAACCATCCTCCGTC 3' (SEQ ID NO. 10)

R: 5' TCATTTTGGCCGACTTAGC3' (SEQ ID NO. 11)

PCR-product size:

108 bp in Col, ca. 104b bp in Ler, (http://genome-www3.stanford.edu/cgi-

BS J

bin/Webdriver?Mlval=atdb\_caps\_max&oid=2f10.210e)

PCR-conditions:

as for med24.2, except that the annealing temperature in the PRC was

raised to 60°C

6. On page 19, the 1<sup>st</sup> full paragraph staring on line 1 should read as follows:

SSLP-marker ng225 (approx. 36 kb proximal of muk11.1 on the sequencing map of

chromosome V; http://www.kazusa.or.jp/arabi/chr5/map/0-2Mb.html)

molecular target:

imperfect (GA)<sub>21</sub> dinucleotide repeat (bases 12203-12244) on

chromosome V BAC-clone MUG13

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PCR primers:

F: 5' TCTCCCCACTAGTTTTGTGTCC 3' (SEQ ID NO. 12)

R: 5' GAAATCCAAATCCCAGAGAGG 3' (SEQ ID NO. 13)

PCR-product size:

119 bp in Col, ca. 189 bp in Ler, (http://genome-www3.stanford.edu/cgi-

bin/Webdriver?Mlval=atdb\_caps\_max&oid=2f10.2112)

PCR-conditions:

as for med24.2, except that MgCl<sub>2</sub> 1.75 mM.

7. On page 19, the 2<sup>nd</sup> full paragraph should read as follows:

<u>CAPS-marker MUG13E</u> (approx. 60 kb proximal of nga225 on the right end of BAC-

clone MUG13 on the sequencing map of chromosome V;

http://www.kazusa.or.jp/arabi/chr5/map/0-2Mb.html)

molecular target:

Accl restriction enzyme site, which is present in the sequence of ecotype

Landsberg erecta, but not Columbia.

PCR primers:

F: 5' GATTTCCCCAGACGATTT 3' (SEQ ID NO. 14)

R: 5' AGTTTATTTGTTGCGGTTTT 3' (SEQ ID NO. 15)

PCR-product size:

2033 bp fragment (bases 79,441-81,473 of BAC MUG13) in Col and Ler

157 ×

PCR-conditions:

before Accl-digest. 1228 and 805 bp fragments in Ler after digest.

as for PAI1-marker, but annealing temperature was 54°C.

Accl-digest:

7µl of PCR-product were mixed with 9µl H20, 2µl 10x buffer M

(Amersham), 2µl BSA (1mg/ml) and 1U Accl, and digested for 5 hrs at

37°C. 10µl of each digest were analyzed on a 1.2% agarose gel.

8. The paragraph starting on the 3<sup>rd</sup> paragraph of page 19 (line 24) and continuing to page 20 line 17 should read as follows:

<u>CAPS-marker PAI2</u> (approx. 33 kb distal of nga158 on the physical map of chromosome V; http://www.kazusa.or.jp/arabi/chr5/pmap/P1\_map 1.html)

molecular target:

Af/III restriction enzyme site, which is present in PAI2-gene from ecotype

Columbia, but not Landsberg erecta.

PCR primers:

F: 5' CAGTTAATGAAACAAGCTTTGTTC 3' (SEQ ID NO. 16)

R: 5' GTTGAGAAAATCACTTTGGTG 3' (SEQ ID NO. 17)

PCR-product size:

645 bp frament (basese 45928-46572 on BAC clone MOP10) in Col and

Ler before Af/III-digest. 590 and 55 bp fragments in Col after digest.

PCR-conditions:

50 mM KCL, 10mM Tris-HCL (pH9.0 @ 25°C), 0.1% Triton? X-100, 250

μM dATP, dGTP, dTTP, dCTP (each), 5 pmoles primer F, 5 pmoles

primer R, 2.5 mM MgCl<sub>2</sub>, 1.0 Units Taq Polymerase (Promega), 40-50 ng

genomic DNA, final volume 25 µl. PCR-program: 1 min 94°C; 35 cycles

(20 sec 94°C, 20 sec 58°C, 90 sec 72°C), 3 min 72°C.

AFIII-digest:

7 μl of PCR-product were mixed with 9μl H20, 2 μl 10 x buffer H

(Amersham), 2µl BSA (1mg/ml) and 1U Af/III, and digested for 5 hours at

37°C. 10 µl of each digest were analyzed on a 2.5% agarose gel.

9. On Page 20, the 2st full paragraph starting on line 18 should read as follows:

<u>SSLP-marker k18i23.1</u> (approx. 101 kb proximal of PAI2 on the sequencing map of chromosome V; <a href="http://www.kazusa.or.jp/arabi/chr5/map/0-2Mb.html">http://www.kazusa.or.jp/arabi/chr5/map/0-2Mb.html</a>)

molecular target:

purine-rich stretch (bases 17830-17870) on chromosome V BAC-clone

K18I123. Length Polymorphism was detected by comparative

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sequencing of the Ler and Col genomic sequence of that region.

PCR primers:

F: 5' TGGTTAGATTTGCTGTT 3' (SEQ ID NO. 18)

R: 5' ATTCTGCATTATTAGTTGTC 3' (SEQ ID NO. 19)

PCR-product size:

139 bp in Col, 133 bp in Ler

PCR-conditions:

as for med24.2, except that MgCl<sub>2</sub> was 2.5 mM and the annealing

temperature during PCR was lowered to 48°C.

10. On Page 21, the first full paragraph starting on line 14 should read as follows:

Genomic DNA for each genotype was prepared from a mixture of young growing leaves and inflorescence tissue using the CTAB-protocol (see above). Six oligonucleotide primers were then designed (Oligo Primer Analysis Software, version 5.0) to amplify three (A,B,C) overlapping PCR-fragments, spanning the entire coding sequence of this cellulose synthase gene on BAC-clone K2A11 (see Figure 2).

B10

Fragment A

was amplified with primers

F<sub>a</sub> (5'-TTAGCCATCCCAAGATTCT-3') (SEQ ID NO. 20) and

R<sub>a</sub> (5'-CTTCAAGGGGTCAACAGTA-3') (SEQ ID NO. 21) giving a

2034 bp PCR-product (bases 13939-15972 on K2A11)

Fragment B

F<sub>b</sub> (5'-TACCGAGCGTTTTTCCTAT-3') (SEQ ID NO. 22)

R<sub>b</sub> (5'-CCAGCACCTAAGTTTCACA-3') (SEQ ID NO. 23)

Fragment C

2064 bp PCR-product (bases 12382-14445 on K2A11)

F<sub>c</sub> (5'-GTTCAGTTCCCACAAAGATT-3') (SEQ ID NO. 24)

R<sub>c</sub> (5'-TCATTCCGACCAAAAGTT-3') (SEQ ID NO. 25)

2395 bp PCR-product (bases 10620-13014 on K2A11)

11. The paragraph starting on the 3<sup>nd</sup> full paragraph of page 23 (line 18) and continuing to page 24 line 4 should read as follows:

The ixr1-2 mutation was mapped with high resolution to a small region on the top arm of chromosome V as described in materials and methods. The mapping results indicated that the ixr1 mutations mapped very near to a gene, Ath-B, encoding cellulose synthase. A cDNA clone for the Ath-B mRNA has previously been described by Arioli et al. (1998). The nucleotidesequence of this clone and the deduced amino acid sequence were deposited in GenBank as accession number AF027174. Comparison of the sequence of the cDNA to the genomic sequence present on BAC clone K2A11 (GenBank accession number AB018111) indicates that the cDNA clone has 87 nucleotides at the 5' end that are not present in the genomic sequence. This extra sequence corresponds to a 59 nucleotide multiple cloning site (G GACTC GCGCGC CTGCAG GTCGAC ACTAGTGGATCC AAA GAATTC G CGGCCG C GTCGAC, (SEQ ID NO. 26) restriction enzymes sites are shown in italics) that was introduced during cloning of the cDNA and an additional 28 nucleotide fragment of DNA (TACGGCTGCGAGAAGACGACAGAAGGGG) (SEQ ID NO. 27) that was also introduced at some stage during the cloning of the cDNA (see bottom insert in Figure 2). A search of GenBank indicated that this sequence is also found at the 5' ends of other cDNA clones; thus it is a common artifact in some libraries.

12. See Sequence Listing attached herewith.

13. See revised Fig. 2 attached herewith.